1	The Menu LINH.	
2	Chairman Phil Mendelson Councilmember Kenyan R. McDuf	fie
3		
4 5		
<i>5</i>		
7	A BILL	
8	ABILL	
9		
10		
11		
12	IN THE COUNCIL OF THE DISTRICT OF COLUMBIA	
13		
14		
15		
16		
17	To amend, on an emergency basis, the Legalization of Marijuana for Medical Treatment	
18	Initiative of 1999 to allow the Alcoholic Beverage and Cannabis Board ("ABC Board")	
19	to issue temporary non-resident registration identification cards that are valid for period	
20	between 3 days and no longer than one year in length, allow licensed testing laboratories	
21	to receive and test samples of medical cannabis products from qualifying patients, allow	V
22 23	licensed testing laboratories to conduct quality assurance or research and development testing for cultivation centers and manufacturers, amend the definition of a social equity	
23 24	applicant to include arrests and convictions of qualifying family members for a cannabi	•
25	or drug offense, expand the list of eligible family members under the social equity	٥.
26	applicant definition to include siblings and grandparents, clarify that existing licensed	
27	cultivation centers and retailers and applicants that scored 150 points or more during th	e
28	open application period that occurred between November 29, 2021 and March 28, 2022	
29	that are authorized by statute to receive a cultivation center, manufacturer, or retailer	_
30	license apart from a designated open application period are not counted in calculating the	ne
31	50% set aside requirement, clarify that the 5 cultivation center registration applicants the	ıat
32	scored 150 points or more during the same open application period shall automatically	
33	receive a manufacturer license provided that they pay the annual fee and register with the	he
34	ABC Board, allow the Alcoholic Beverage and Cannabis Board to issue conditional	
35	licenses to testing laboratory applicants, and to waive the application fee for testing	
36	laboratory licenses.	
37		
38	BE IT ENACTED BY THE COUNCIL OF THE DISTRICT OF COLUMBIA, That the	IS
39	act may be cited as the "Medical Cannabis Clarification and Non-Resident Patient Access	

Emergency Amendment Act of 2023".

41	Sec. 2. The Legalization of Marijuana for Medical Treatment Initiative of 1999, effective
42	February 25, 2010 (D.C. Law 13-315; D.C. Official Code § 7-1671.01 et seq.), is amended as
43	follows:
44	(a) Section 2 (D.C. Official Code § 7-1671.01) is amended as follows:
45	(1) Paragraph (13B)(B) is amended by striking the phrase "30-day registration
46	identification card" and inserting the phrase "registration identification card valid for periods
47	established by the ABC Board by rulemaking, which are between 3 days and no longer than one
48	year in length" in its place.
49	(2) Paragraph (20C)(B) is amended by striking the phrase "or has a non-parent legal
50	guardian who is or has been incarcerated" and inserting the phrase "or has a non-parent legal
51	guardian, or a grandparent or a sibling who is or has been arrested, convicted, or incarcerated".
52	(b) Section 6(b) (D.C. Official Code § 7-1671.05(b)) is amended as follows:
53	(1) Paragraph (4) is amended as follows:
54	(A) Subparagraph (A) is amended by striking the phrase "30 days" and
55	inserting the phrase "periods established by the ABC Board by rulemaking, which are between 3
56	days and no longer than one year in length".
57	(B) Subparagraph (B) is amended by striking the phrase "30-day".
58	(2) Paragraph (5)(C) is amended by striking the phrase "3 years." and inserting the
59	phrase "3 years, except for temporary non-resident registration identification cards that are valid
60	for periods established by the ABC Board by rulemaking, which shall be between 3 days and no
61	longer than one year in length." in its place.
62	(3) A new paragraph (11A) is added to read as follows:
63	"(11 A) Allow testing laboratories to:

"(A) Receive and test samples of medical cannabis products from
qualifying patients; provided, that the qualifying patient must present proof that he or she is
currently registered, and that the medical cannabis product was purchased from a retailer or
internet retailer licensed with ABCA.

- "(B) Receive and test samples of medical cannabis products from licensed cultivation centers or manufacturers for purposes of quality assurance or research and development. Samples collected for quality assurance or research and development testing may be selected by the cultivation center or manufacturer non-randomly. Any tests conducted for purposes of quality assurance or research and development shall not satisfy the requirements of paragraphs (8) through (11) of this subsection."
 - (c) Section 7 (D.C. Official Code § 7-1671.06) is amended as follows:
- (1) Subsection (h) is amended by striking the phrase "cultivation centers who receive a manufacturer's license pursuant to subsection (d) of this section" and inserting the phrase "cultivation centers and retailers, and applicants who scored 150 points or more during the ABC Board open application period that occurred between November 29, 2021 and March 28, 2022, who receive a cultivation center, manufacturer, or retailer's license pursuant to subsections (d), (w), (x) and (y) of this section."
 - (2) Subsection (k)(1) is amended to read as follows:
- "(k)(1) The ABC Board shall be authorized to issue a one-year conditional license for a cultivation center, retailer, internet retailer, manufacturer, courier, or testing laboratory that does not currently have a proposed location.".
 - (3) Subsection (n)(2) is amended to read as follows:

86	"(2)(A) The ABC Board shall, by rules issued pursuant to section 14, establish the
87	initial application and renewal fees for cultivation center, manufacturer, retailer, internet retailer,
88	and courier licenses. The ABC Board may revise these fees as considered necessary.
89	"(B) There shall be no initial application fee for a testing laboratory
90	license. Renewal fees for a testing laboratory license shall be established by rules issued pursuant
91	to sub-paragraph (A) of this paragraph.".
92	(3) Subsection (y) is amended as follows:
93	(A) Strike the phrase "be considered by the ABC Board for a manufacturer
94	license" and insert the phrase "automatically receive a manufacturer license provided that the
95	annual fee is paid.
96	(B) Strike the phrase "files a manufacturer license application" and insert
97	the phrase "registers on a form provided by ABCA" in its place.
98	Sec. 3. Repealer.
99	The Medical Cannabis Manufacturer Clarification Temporary Amendment Act of 2023,
100	passed on 2nd reading on June 20, 2023 (Enrolled version of Bill 25-304), is repealed.
101	Sec. 4. Fiscal impact statement.
102	The Council adopts the fiscal impact statement of the Budget Director as the fiscal impact
103	statement required by section 4a of the General Legislative Procedures Act of 1975, approved
104	October 16, 2006 (120 Stat. 2038; D.C. Official Code § 1-301.47a).
105	Sec. 5. Effective date.
106	This act shall take effect following approval by the Mayor (or in the event of veto by the
107	Mayor, action by the Council to override the veto), and shall remain in effect for no longer than
108	90 days, as provided for emergency acts of the Council of the District of Columbia in section

- 109 412(a) of the District of Columbia Home Rule Act, approved December 24, 1973 (87 Stat. 788;
- 110 D.C. Official Code § 1-204.12(a)).